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COMPLETE SPECIFICATION

Improvements relating to Halogen Substituted Diphenyl Urea and Thiourea Compounds and their use

We, J. R. GEIST A.—G., a body corporate organised according to the laws of Switzerland, of 215 Schwarzwaldallee, Basle, Switzerland, do hereby declare the invention for 5 which we pray that a patent may be granted to us, and the method by which it is to be performed, to be particularly described in and by the following statement:—

The present invention concerns the production of polyhalogen substituted monohydroxydiphenyl urea and thourea compounds which have at least one halogen substituent in each of the two benzene rings but no acid water solubilising groups. It also concerns their use

15 as disinfectants.

o-Hydroxydiphenyl urea and thiourea compounds which have at least one halogen substituent in each of the two benzene rings but no acid water solubilising groups, are new. They
can be produced by methods known per se by reacting halogen substituted o-hydroxyaminobenzene compounds with a compound introducing a halogen phenyl carbamyl- or thiocarbamyl radical into the primary amino group. 25 Chiefly the halogen substituted phenyl isocyanates and/or phenyl mustard oils can be used as compounds which introduce the halogen phenyl carbamyl or thiocarbamyl radical. Phenyl carbamic acid phenol esters which are asily obtained from chloroformic acid phenol esters and halogen aminobenzenes can also be used as starting materials in the process according to the present invention, as, at even moderate temperatures with correspondingly 35 chosen halogen-o-hydroxyaminobenzene compounds, they produce o-hydroxydiphenyl ureas according to the present invention whilst splitting off phenol. Also the reaction of suitably substituted phenyl ureido compounds with o-40 aminohydroxybenzene compounds chosen according to this invention leads, in individual cases, to polyhalogen substituted o-hydroxy-diphenyl ureas. Finally, also the use of halogen

benzovlazides, which under the conditions of

[Price 3s. Od.]

the reaction known per se transform themselves whilst splitting off intogen into the corresponding halogen phenyl isocyanates, falls within the scope of the present invention. However, the addition of balogen phenyl isocyanate or halogen phenyl insurant oil to suitably substituted halogen-neuminobenzene compounds is to be preferred to all other methods.

Examples of phenyl isocyanate or habut

Examples of phenyl isocyanates or phenyl mustard oils which can be used in the process according to the present invention are: 4chlorophenyl isocyanate, 3.4-dichlorophenyl isocyanate, 3-trifluoromethyl-4-chlorophenyl isocyanate, 3-4-bromophenyl isocyanate, 4-fluorophenyl isocyanate, 3-4-dibromophenyl isocyanate, 3-4-tirichlorophenyl isocyanate, 2-3-4-trichlorophenyl isocyanate, 4-fluoro - 3chlorophenyl isocyanate, 4-chlorophenyl mustard oil, 3.4-dichlorophenyl mustard oil, 3 trifluoromethyl-4-chlorophenyl mustard oil. They are reacted according to the invention with halogen substituted o-aminophenols. Examples of such are: 4-chloro-2-amino-1hydroxybenzene, 5 - chloro - 2 - amino - 1 hydroxybenzene, 4.5 - dichloro - 2 - amino - 1- 70 hydroxybenzene, 3.4.6 - trichloro - 2 - amino-1 - hydroxybenzene, 4- or 5-bromo-2-amino-1-hydroxybenzene, 4-6-dibromo- or 4-6-dichloro - 2 - amino - 1 - hydroxybenzene, 4.5dibromo - 2 - amino - 1 - hydroxybenzene, 75 4 - chloro - 5 - trifluoromethyl-2-amino - 1-hydroxybenzene, 4 - chloro - 5 - bromo - 2 -

aminophenyl.

Phenyl carbamic acid phenol esters usable in the process according to the present inven-80 tion are: 3.4- dichlorophenyl carbamic acid phenol ester or o-cresyl ester, 3-trifluoro-methyl -4- chlorophenyl carbamic acid phenol or o-creyl ester, 4-tromophenyl carbamic acid phenol ester 3.4-dibromophenyl carbamic acid 85 phenol ester 3.4-dibromophenyl curbamic acid 85 phenol ester 3.4-dibromophenyl curbamic acid

According to the present invention the com-

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ponents are so chosen that there is at least one halogen substituent in each benzene ring in the new 2-hydroxydiphenyl urea or thiourea compounds whereby in the benzene ring containing no hydroxyl group, the halogen substituent can be replaced by the trifluoromethyl group.

It is preferable that these halogen substituents occupy the p- and also, if desired, the m-position to the urea or thiourea bridge. Halogen substitution in pairs in the m- and p-positions to the ureido group in at least one of the benzene rings is particularly advantageous, If desired, allyl or alkoxy groups can be present as further substituents, e.g. a methyl group in either the m- or p-position to the hydroxyl group or in the c-position to the treat or and all the control of the

hydroxyl group.
Particularly valuable o-hydroxydiphenyl ureas according to this invention correspond to the general formula:

$$x_1 \rightleftharpoons NH - C - NH - C X_4$$

wherein X1 to X4 represent halogen or hydro-

X₂ in addition may represent the trifluoromethyl group whilst not more than one of the four symbols X represents hydrogen, and Z represents an oxygen or a sulphur atom, and wherein the two bearzene rings can each contri

and wherein the two benzene rings can each contain a further substituent, e.g. another halogen or a methyl group. Some of the polyhalogen substituted ohydroxydiphenyl urea or thiourea compounds,

35 with suitable halogen substitution, have a very good bactericidal action. In particular they are very active against the bacterial flora which cause perspiration odours and they are, for this season and because of their low rozincity, very suitable for use as decodorants in laundering, and for human use when incorporated in cleansing agents such as soaps or shampoos, or as additives to cosmetic agents such as sointments or creams.

The following examples illustrate the invention without limiting it in any way. Where not otherwise stated, parts are given as parts by weight and the temperatures are in degree Centigrade. The relationship of parts by weight to parts by volume is as that of kiloeramnes to litres.

Example 1.

A hot solution of 21.5 parts of 3.4-dichlorobenzoic acid azide in 50 parts by volume of follone is added to a hot solution of 14.5 parts of 5-chloro-2-aminophenol in 250 parts by volume of roluten. After boiling for 15 minutes and then cooling the mixture, the N-3.4 dichlorophenyl-N-2-lydroxy - 4* - chloro - phenyl urea formed is filtered off. When recrystallised from diluted alcohol, it melts at 201—202°. This compound also has desirable fungicidal properties.

EXAMPLE 2.

A mixture of 22.5 parts of 3.4-dichloro-phenyl urea, 14 parts of 2-amino-4-chloro-phenol and 40 parts by volume of glacial accided is boiled until no more 2-amino-4-chloro-phenol cam be traced. The cooled mass is diluted with a little water whereupon it solidifies. The precipitate is filtered off and extracted with diluted caustic soda to remove the 5-chloroberoxeosione. Lower of the cooled with the addition of caustic soda by earlies and a lower of the filtered off and the N-3.4-dichlorophenyl N2-lydroxy5-chlorophenyl urea is precipitated with acetic acid. After recrystallising from disoane it melts at 203–206°.

EXAMPLE 3.

31 Parts of 3.4-dichlorophenyl carbamic acid phenyl ester and 17 parts of 2-amino-4.5-dichlorophenol are dissolved in 100 parts by volume of dioxane and 50 parts of a 40% sodium acctate solution are added. The mixture is stirred at 90—95' until for practical purposes no more 2-amino-4.5-dichlorophenyl and 100 parts of 100 parts of 100 purposes no more 2-amino-4.5-dichlorophenyl are which has partly precipitated during the reaction is fiftered off after cooling and recrystallised from tetrachlore-ethane. It melts at 201—202.

EXAMPLE 4.

18 Parts of 2-amino-4.5-dichlorophenol are 94 dissolved in 25 parts by volume of acctone and a solution of 17 parts of 4-chlorophenyl mustard oil in 25 parts by volume of acctone is added. The whole is stirred for 3 hours at 35, diluted with water and he N-4-chlorophenyl - N'.2' - hydroxy - 4'.5' - dichlorophenyl thiourea which precipitates is filtered off. After recrystallising from benzene it melts at 146—147' on decomposition

EXAMPLE 5.

A solution of 10 parts of 3.4-dichlorophenyl mustred oil in 40 parts by volume of benzene is added to a solution of 10 parts of 2-amino-4.5-dichlorophenol in 20 parts py volume of acetone. After stirring for 3 hours, the product is precipitated with pertoleum ether and the separated N-3.4-dichlorophenyl - N'.2' hydroxy - 4'5' - dichlorophenyl thiourea is recrystallised from benzene, M.P. 159—160' on decomposition.

Example 6.

100 Parts of an 18% solution of 3.4-dichlorophenyl isocyanate in nitrobenzene are added at 30—35° while cooling to a solution of 19 parts of 2-amino-5-bromophenol in 25 parts by volume of acctone. The whole is stirred for an hour, filtered, the product is washed with benzene and recrystallised from diluted alcohol. The N-3-4-dichlorophenyl-N'-2-hydroxy-4'-bromophenyl urea obtained melts at 198—199°.

EXAMPLE 7.

100 Parts of an 1836, solution of 3.4-di10 chlorophenyl isocynate in nitrobenzue are
poured at 30—35° into a solution of 23 parts of
2 amino-4-chloro-5-bromophenol in 40
parts by volume of acetone and the whole is
streed for 1 hour. The N-3.4- delherophenyl15 N'2-1-ydroxy-4-bromo-5'-chlorophenyl urea
is filtered off, boiled with benzene to remove
the nitrobenzuene, again filtered off and recrystallised from diluted alcohol, M.P. 201—
202° on decomposition.

EXAMPLE 8.

111 Parts of a 9% solution of 3-triflacormethyl-4-chilorophenyl isocepante in chloropbenzene is poured at 30° into a solution of 9
parts of 2-amino-4-3-chilorophenol in 25 parts
by volume of actome. After stirring for 2 hours,
the product is filtered off and recrystallised
from benzene. N-3-triflacomethyl -4-chilorophenyl-N-22 - hydrawy - 4-3-dichlorophenyl
urea melts at 174—175°.

EXAMPLE 9.

123 Parts of a 9% solution of 3-trifluoromethyl-4-chlorophenyl isocynaete are poured at 30—35° into a solution of 11 parts of 3.46-trichloro-2-minophenol in 30 parts by volume of accume. On completion of the reaction, the N - 3 - trifluoromethyl-4-chlorophenyl - N;2 - hydroxy-3;5:6° trichlorophenyl urea formed is filtered off and recrystallised from diluted alcohol, MP, 198—199°.

Example 10.

8 parts of 2-amino-4.6-dichlorophenyl are dissolved in 70 parts by volume of acetone and the solution is poured at room temperature into 8 parts of 2.3.4-trichlorophenyl isocyanate in 30 parts by volume of acetone. After stirring for 2 hours, the product is filtered off. Recrystallised from glacial acetic acid, N-2.3-4trichlorophenyl - N'.2' - hydroxy - 3'.5' dichlorophenyl urea melts at 214-215.

EXAMPLE 11.

100 Parts of an 18½ solution of 3.4 - dichitorophenyl isocynate in introhenzene are added at 30° to a solution of 22 parts of 2amino-4.5.6-tichlorophenol in 50 parts by volume of acetone. After stirring for one hour, the N-3.4-dichtorophenyl - N'22 - hydroxy-3'.4'5'-trichlorophenyl urea is precipitated with petroleum ether, filtered off and recrystallisted from chlorobenzene. M.P. 210—211° on decomposition

The bactericidal properties and the melting points of some diphenyl urea or thiourea compounds according to this invention can be sen from the following Table, that gives puriticlars from which said compounds can be identified. The bactericidal properties were determined on Staphylococcus aureus as follows:

A standard suspension, which is prepared by adding sterilised tap water to the germs of 16 hour agar cultures, the density of which is brought to 85% transparency in the so-called BIO-PHOTO-COL-apparatus according to Hellige, is mixed with graduated dilutions of the disinfectant to be tested (in aqueous solution). Duration of test: 10 minutes, temperature: 20°.

At the end of the 10 minutes, 2 sub-cultures from each reaction mixture are prepared with a glucose broth. The sub-cultures are bred at 37° C. After 48 hours, the development of sterility of the sub-cultures is determined. The bactericidal activity of a disinfectant is determined by the minimum concentration required to full, with certainty, a standard supersion of test germs under certain conditions. The minimum is accretioned by graduated consolutations according to the dilution process principle and is expressed in 10°4 mol.

TABLE

o-aminophenol	phenyl	M.P.	minimum bactericidal concentration expressed in 10 ⁻⁶ mol
CI OH	≺Ci ci	206—207°	25
CI-Q-WH	-⇔cı	201—202°	12.5
31-0-NH-	-⇔a	198—199°	25
OH -	ci ci	204—205°	25
Br - NH-	~ €0	201—202° on decomposition	12.5
CT OH WH-	-⇔cı c _ξ	174—175°	6.2
CL-OH-	- de er	201—202	3.1
CI -NH -	****	214—215°	12.5
CI - NH - CI CI	-√ar	198—199°	6.2
	OR NH- OH OH NH- OH NH-	OH NH- OH O	OH

No.	o-aminophenol	phenyl	M.P.	minimum bactericidal concentration expressed in 10 ⁻⁶ mol
10.	CI NH-	₹a	210—211° on decomposition	. 3.1
11.	CI - NH -	Thioureas:	146—147° on decomposition	12.5
12.	CI CI NH-	⇔a a	159—160° on decomposition	12.5

EXAMPLE 12.

99 Parts of soap fiekes and I part of N-3-ddichloropheny! N-72 - hydroxy - 45;65 chlorophenyl urea together with a little perfume are well mixed either direct or dissolved in a little alcohol in a mixing apparatus. The finished mixture is refined by rolling and then pressed into tablet form. A toilet soap having 10 a deodorant action is thus obtained.

A similar good action is obtained if 97 parts of soap flakes and 3 parts of N-3.4-dichlorophenyl - N'.2' - hydroxy - 3'.4'.5'-trichlorophenyl urea are used as starting materials.

Example 13.

A 1% aqueous solution of a cleansing agent which contains 10 parts of one of the diphenyl ureas named in example 12 and 90 parts of a non-ionogenic or anion active synthetic washing agent, produces a washing ilquor which is suitable for example for the cleansing of household or personal linen.

What we claim is:—

1. A polyhalogen substituted compound hav-

25 ing the general formula:

wherein Hal represents halogen,

Y represents halogen or the trifluoromethyl group, and

Z represents an oxygen or sulphur atom; and the two benzene rings may contain further substituents but no water solubilising groups. 2. A polyhalogen substituted compound

A polyhalogen substituted compound having the formula:

wherein X1, X2, X3 and X4 represent hydrogen or halogen,

Y represents halogen or the trifluoromethyl

Z represents an oxygen or sulphur atom. 4
3. A polyhalogen substituted compound having the formula:

wherein X₁, X₂, X₃ and X₄ represent halogen or hydrogen,

X₃ may also represent the trifluoromethyl group whilst not more than one of the four symbols X represents hydrogen, and

Z represents an oxygen or a sulphur atom.

4. Process for the production of polyhalogen 50, substituted monohydroxydiphenyl urea or thiourea compounds, the benzene rings of which contain no acid water solubilising groups, characterised by reacting halogen substituted o-hydroxyaminobenzene compounds 55 with a compound which introduces a halogen or trifluoromethyl substituted phenyl carbamyl crificial into the nrimary

amino group.

5. Manufacture of polyhalogen substituted compounds substantially as herein described

with reference to any of the foregoing examples 1 to 11. A polyhalogen substituted compound as hereinbefore identified by or in any of the examples 1 to 11 or by the Table.

7. Germicidal agent characterised by a con-

tent of a compound of the general formula:

wherein Hal represents halogen, Y represents halogen or the trifluoromethyl

Z represents an oxygen or a sulphur atom, and the two benzene rings may contain further substituents but no water solubilising groups.

8. A process of forming a toilet soap containing a polyhalogen substituted compound, substantially as described in Example 12. 9. A process of producing a liquor suitable

A pacces or producing a liquor suitable for cleansing household or personal linen and containing a polyhalogen substituted com-pound, substantially as described in Example 13.

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